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IN THE CLAIMS

Please amend the claims as follows.

1. (Currently amended) A method of analyzing a sample for the presence of a member of a specific binding pair, the method comprising:

providing a microsphere having an incorporated electroactive marker encapsulated within the microsphere and a first member of a specific binding pair attached to the microsphere wherein the microsphere is not a liposome;

introducing a sample suspected to comprise a second element of the specific binding pair complex to the miscrosphere;

selecting for the microsphere by formation of a specific binding pair complex <u>in fluid</u> <u>suspension</u>; and

detecting the specific binding pair complex by electrochemical testing for the electroactive marker released from the microsphere

wherein,

electrochemical testing is via voltammetry or amperometry.

- 2. (Original) The method of claim 1 wherein the microsphere is a polymeric microsphere that is insoluble in an aqueous solution.
- 3. (Original) The method of claim 2 wherein the microsphere is a polystyrene-based microsphere.
 - 4. (Cancelled)
- 5. (Currently amended) The method of claim 4-1, wherein providing comprises incubation of a polymeric microsphere in an organic solvent including an the electroactive marker.

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6. (Cancelled)

7. (Previously presented) The method of claim 1 wherein selecting comprises binding

of the first member of the specific binding pair attached to the microsphere and a second member

of the specific binding pair attached to a substrate.

8. (Original) The method of claim 7 wherein the first member of the specific binding

pair attached to the microsphere comprises a covalent bond with a functional group on the

surface of the microsphere.

9. (Original) The method of claim 7 wherein the substrate comprises a magnetic particle.

10. (Original) The method of claim 1 wherein selecting comprises incubation.

11. (Original) The method of claim 1 wherein the specific binding pair complex is an

antigen/antibody, enzyme/substrate, oligonucleotide/DNA, chelator/metal, enzyme/inhibitor,

bacteria/receptor, virus/receptor, hormone/receptor, DNA/RNA, RNA/RNA, or

oligonucleotide/RNA complex.

12. (Cancelled)

13. (Currently Amended) The method of claim 1 wherein releasing released comprises

solubilizing the microsphere.

14. (Original) The method of claim 1 wherein the electroactive marker comprises a

metallocene.

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15. (Original) The method of claim 1 wherein the electroactive marker comprises a

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nanoparticle.

16. (Original) The method of claim 1 wherein the electroactive marker comprises a

metal.

17. (Original) The method of claim 1 wherein electrochemically testing comprises

measurement of one or more electrical quantities in relationship to one or more chemical

parameters.

18. (Currently amended) The method of claim 14 17 wherein the electrical quantities

comprises current, potential or charge.

19. (Cancelled)

20. (Currently amended) A method of analyzing a sample for the presence of two or

more analytes, the method comprising:

providing a first microsphere having an incorporated a first electroactive marker

incorporated into the body of the first microsphere;

providing a second microsphere having a incorporated second electroactive marker

electrochemically distinguishable from the first electroactive marker encapsulated within the

body of the second microsphere wherein neither the first microsphere nor the second

micropshere is a liposome;

attaching a first binding pair member specific to a first analyte to the first microsphere;

attaching a second binding pair member specific to a second analyte to the second

microsphere;

incubating the first microsphere and second microsphere in a solution comprising the

sample to be analyzed;

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selecting for the first microsphere and second microsphere by formation of specific binding pair complexes <u>in fluid suspension</u>; and

detecting the specific binding pair with electrochemical testing for the first electroactive marker and the second electroactive marker <u>released from the microsphere</u>

wherein,

electrochemically detection is via voltammetry or amperometry

- 21. (Original) The method of claim 20 wherein the microsphere is a polymeric microsphere that is insoluble in an aqueous solution.
- 22. (Original) The method of claim 21 wherein the microsphere is a polystyrene-based microsphere.
- 23. (Original) The method of claim 20 wherein attaching comprises a covalent bond with a functional group on the surface of the microsphere.
- 24. (Original) The method of claim 20 wherein the specific binding pair complexes are antigen/antibody, enzyme/substrate, oligonucleotide/DNA, chelator/metal, enzyme/inhibitor, bacteria/receptor, virus/receptor, hormone/receptor, DNA/RNA, RNA/RNA, or oligonucleotide/RNA complexes.
- 25. (Original) The method of claim 20 further comprising releasing the first electroactive marker from the first microsphere and the second electroactive marker from the second microsphere.
- 26. (Currently amended) The method of claim 25 20 wherein releasing comprises solubilizing the first microsphere and the second microsphere.

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27. (Original) The method of claim 20 wherein the first electroactive marker and the

second electroactive marker comprise metallocenes.

28. (Original) The method of claim 20 wherein the first electroactive marker and the

second electroactive marker comprise nanoparticles.

29. (Original) The method of claim 20 wherein the first electroactive marker and the

second electroactive marker comprise metal.

30. (Original) The method of claim 20 wherein electrochemically testing comprises

measurement of one or more electrical quantities in relationship to one or more chemical

parameters.

31. (Original) The method of claim 30 wherein the electrical quantities comprises

current, potential or charge.

32. (Cancelled)

33. (Withdrawn) A microsphere for electrochemical detection of a member of a specific

binding pair, comprising a polymeric microsphere having an organic solvent soluble

hydrophobic electroactive marker incorporated into the body of the microsphere and at least one

functional group on the surface of the microsphere.

34. (Withdrawn) The microsphere of claim 33 wherein the soluble hydrophobic

electroactive marker is non-magnetic.

35. (Withdrawn) The microsphere of claim 34 wherein the soluble hydrophobic

electroactive marker is a metallocene.

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36. (Withdrawn) The microsphere of claim 35 wherein the metallocene is ferrocene or

ferrocenecarboxaldehyde.

37. (Withdrawn) The microsphere of claim 33 wherein the at least one functional group

is a sulfate surface group, aldehyde group, aliphatic amine group, amide group, aromatic amine

group, carboxylic acid group, chloromethyl group, epoxy group, hydrazide group, hydroxyl

group, sulfonate group or tosyl group.

38. (Withdrawn) The microsphere of claim 33 wherein the polymeric microsphere is a

polystyrene-based microsphere.

39. (Withdrawn) The microsphere of claim 38 wherein the polystyrene-based

microsphere has a diameter between about 0.01 μm and about 100.0 μm.

40. (Withdrawn) The microsphere of claim 38 wherein the polystyrene-based

microsphere has a diameter between about 0.3 μm and about 20 μm .